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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
10/672,484	09/25/2003	Roland Contreras	13748Z	8325	
2389 7590 09/16/2008 SCULLY SCOTT MURPHY & PRESSER, PC 400 GARDEN CITY PLAZA			EXAM	EXAMINER	
			NGUYEN, QUANG		
SUITE 300 GARDEN CIT	Y. NY 11530	ART UNIT	PAPER NUMBER		
	-,		1633		
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Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Office Action Summary

Application No.	Applicant(s)	
10/672,484	CONTRERAS ET AL.	
Examiner	Art Unit	
QUANG NGUYEN, Ph.D.	1633	

Period fo	The MAILING DATE of this communication appears on the cover sheet with the correspondence address or Reply				
WHIC - Exte after - If NC - Failu Any	ORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE 2 MONTH(S) OR THIRTY (30) DAYS, CHEVER IS LONGER, FROM THE MAILING DATE OF THIS COMMUNICATION. Insorts of time may be available under the provisions of 37 CFR 1.138(a). In no event, however, may a reply be timely filed SIX (5) MONTHS from the mailing date of this communication. The provision of time may be available under the provisions of 37 CFR 1.138(a). In no event, however, may a reply be timely filed SIX (5) MONTHS from the mailing date of this communication. The provision of the set or extended period for reply will the platable, cause the application to become ARAMONDED (SU SC. \$1.33). The provision of the provisi				
Status					
1)🖂	Responsive to communication(s) filed on 13 June 2008.				
	This action is FINAL. 2b) This action is non-final.				
3)	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213.				
Disposit	ion of Claims				
4)⊠	Claim(s) 74-113 is/are pending in the application.				
,	4a) Of the above claim(s) 74-89 is/are withdrawn from consideration.				
5)	Claim(s) is/are allowed.				
6)🖂	Claim(s) <u>90-113</u> is/are rejected.				
	Claim(s) is/are objected to.				
8)□	Claim(s) are subject to restriction and/or election requirement.				
Applicat	ion Papers				
9)[The specification is objected to by the Examiner.				
10)	The drawing(s) filed on is/are: a) ☐ accepted or b) ☐ objected to by the Examiner.				
	Applicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a).				
	Replacement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d).				
11)	The oath or declaration is objected to by the Examiner. Note the attached Office Action or form PTO-152.				
Priority (ınder 35 U.S.C. § 119				
	Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). ☐ All b □ Some * c) ☐ None of:				
	 Certified copies of the priority documents have been received. 				
	2. Certified copies of the priority documents have been received in Application No				
	 Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). 				
* 5	See the attached detailed Office action for a list of the certified copies not received.				
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Attachmen	(fe)				
_	te of References Cited (PTO-892) 4) Interview Summary (PTO-413)				

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Attachment(s)		
Notice of References Cited (PTO-892)	4) Interview Summary (PTO-413)	
Notice of Draftsperson's Patent Drawing Review (PTO-948)	Paper No(s)/Mail Date	
3) Tinformation Disclosure Statement(s) (PTO/SS/08)	 Notice of Informal Patent Application 	
Paper No(s)/Mail Date	6) Other:	

DETAILED ACTION

Applicant's election with traverse of the following species in the reply filed on 6/13/08 is acknowledged. Applicants elected: (a) fungal glucosidase II and (b) GAP promoter.

Upon further consideration and in light of the prior art rejections applied below, the species restriction requirement is withdrawn.

Claims 74-113 are pending in the present application.

This application contains claims 74-89 drawn to an invention nonelected with traverse in the reply filed on 8/9/06. A complete reply to the final rejection must include cancellation of nonelected claims or other appropriate action (37 CFR 1.144) See MPEP § 821.01.

Accordingly new claims 90-113 are examined on the merits herein.

Response to Amendment

The rejection under 35 U.S.C. 112, first paragraph, for the Lack of Written Description was withdrawn in light of Applicant's amendment.

The rejection under 35 U.S.C.102(b) as being anticipated by Martinet et al. (Biotechnology Letters 20:1171-1177, 1998; IDS) as evidenced by the pPICZB vector diagram (Invitrogen Catalog, 1998; IDS) was withdrawn in light of Applicant's amendment.

The rejection under 35 U.S.C. 102(b) as being anticipated by Chiba et al. (J. Biol. Chem. 41:26298-26304, 1998; IDS) as evidenced by Inoue et al. (Biochim. Biophys. Acta 1253:141-145, 1995; IDS) was withdrawn in light of Applicant's amendment.

The rejection under 35 U.S.C. 102(b) as being anticipated by JP 8-336387 (IDS) was withdrawn in light of Applicant's amendment.

Claim Rejections - 35 USC § 112

The following is a quotation of the second paragraph of 35 U.S.C. 112:

The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.

New claims 107-113 are rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention. This is a new ground of rejection necessitated by Applicant's amendment.

New claims 107-113 are rejected under 35 U.S.C. 112, second paragraph, as being incomplete for omitting essential elements, such omission amounting to a gap between the elements. See MPEP § 2172.01. The omitted element is: the *Pichia* strain must also be transformed with a nucleotide sequence coding for a heterologous glycoprotein so that the transformed cells of said *Pichia* strain can produce the heterologous glycoprotein. As written, there is no linkage between the preamble of the claims reciting "reducing glycosylation of a heterologous glycoprotein" with the body of the claim simply reciting transforming cells of a *Pichia* strain with a nucleotide sequence coding for a *T. reesei* a-1,2-mannosidase or its functional part thereof and a nucleotide

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sequence comprising a portion of the genomic OCH1 gene of said strain operably linked to a selectable marker

Additionally, independent claim 107 recites the limitation "said functional part thereof" in lines 3 and 5 of the claim. There is insufficient antecedent basis for this limitation in the claim. This is because prior to these limitations, there is no recitation of functional part of anything. Accordingly, the metes and bounds of the claims are not clearly determined.

Claim Rejections - 35 USC § 102

The following is a quotation of the appropriate paragraphs of 35 U.S.C. 102 that form the basis for the rejections under this section made in this Office action:

A person shall be entitled to a patent unless -

(e) the invention was described in (1) an application for patent, published under section 122(b), by another filed in the United States before the invention by the applicant for patent or (2) a patent granted on an application for patent by another filed in the United States before the invention by the applicant for patent, except that an international application filed under the treaty defined in section 351(a) shall have the effects for purposes of this subsection of an application filed in the United States only if the international application designated the United States and was published under Article 21(2) of such treaty in the English language.

New claims 90-96 and 104-113 are rejected under 35 U.S.C. 102(e) as being anticipated by Gerngross (US 2002/0137134; IDS) as evidenced by JP 8-336387 (IDS).

This is a modified rejection necessitated by Applicant's amendment.

Gemgross discloses methods and compositions by which fungi or other eukaryotic microorganisms including <u>Pichia pastoris</u>, <u>Hansenula polymorpha</u>, <u>Candida albicans</u> can be genetically modified to produce glycosylated proteins having patterns of glycosylation similar to glycoproteins produced by animal cells, particularly human cells,

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which are useful as human or animal therapeutic agents such as erythropoietin, cytokines, coagulation factors (See Summary of Invention, pages 4-5; page 6. paragraph 56). This is achieved using a combination of engineering and/or selection of strains which do not express a certain enzymes that which create the undesirable complex structures characteristic of the fungal glycoproteins, which express exogenous enzymes selected either to have optimal activity under the conditions present in the fungi where activity is desired or which are targeted to an organelle where optimal activity is achieved (including α-1,2-mannosidase from Trichoderma reesei, paragraph 68-71), and combinations thereof wherein the genetically engineered eukaryote expresses multiple exogenous enzymes required to produce "human-like" glycoproteins (see abstract; paragraphs 41, 48). Specifically, Gerngross teaches that in a preferred process or embodiment the microorganism is engineered to express an exogenous α-1,2-mannosidase enzyme having an optimal pH between 5.1 and 8.0, and that the enzyme is targeted the endoplasmic reticulum (ER) or Golgi apparatus of the host organism, where it trims N-glycans such as ManaGlcNAc2 to yield Man₅GlcNAc₂ which is a substrate for further glycosylation reactions that produce a finished N-glycan that is similar or identical to that formed in mammals and it is not a substrate for hypermannosylation reactions that occur in vivo in yeast or other microorganisms (paragraphs 42, 68-72; Table 6 and claim 1). Gerngross also teaches that ER or Golqi apparatus targeting sequences are well known in the art such as HDEL or KDEL (paragraphs 87-88 and Table 6); and where the host is Pichia pastoris suitable promoters such as AOX1, AOX2, DAS and P40 promoters can be used at least for Application/Control Number: 10/672,484 Page 6

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expression of an exogenous gene encoding glycosylation enzymes (paragraph 85 and example 1). Gerngross specifically teaches that the eukaryotic strains which do not express one or more enzymes involved in the production of high mannose structures are used, and that these strains can be engineered in conjunction with the introduction of an exogenous α -1,2-mannosidase enzyme (paragraphs 64-67), and one of the many such mutants already described in yeasts including a hypermannosylation-minus (OCH1) mutant in Pichia pastoris described in Japanese Patent Application Public No. 8-336387 (paragraphs 35, 48). JP 8-336387 already teaches the preparation of a vector construct comprising a portion of Pichia OCH1 gene and a selectable marker gene for disruption of the genomic OCH1 in a Pichia yeast strain, including the GTS 115 (NRRL Y-15851) strain for inhibiting the elongation of sugar chains on glycoproteins (see abstract in English, Fig. 9 on page 635 as well as col. 15, paragraph 0033). It is further noted that articles (e.g., a cultured flask or a vial) containing vectors for expression of exogenous α-1,2-mannosidase enzyme and for disruption of the genomic OCH1 in a Pichia yeast strain and/or the eukaryotic microorgainisms (e.g., Pichia veasts) in the invention of Gerngross would constitute such a kit.

Accordingly, the teachings of Gerngross meet every limitation of the instant claims. Therefore, the reference anticipates the instant claims.

Response to Arguments

Applicants' arguments related to the above rejection in the Amendment filed on 5/3/07 (pages 15-17) have been fully considered but they are respectfully not found persuasive for the reasons discussed below.

1. Applicants argue that Gerngross does not teach the strains and methods as presently claimed, particularly *Pichia pastoris* that is transformed with a vector capable of expressing a *T. reesei* α-1,2-mannosidase or a functional part thereof. Instead, Gerngross teaches away from such a strain because Gerngross discloses that to obtain Man₅GlcNAc₂ in high yield, one could engineer a strain that expresses a α-1,2-mannosidase which should have an optimal pH that is between 5.1 and 8.0, while the *T. reesei* α-1,2-mannosidase has an optimal pH of 5.0 and Gerngross discusses the inefficiency of this enzyme in several passages.

Firstly, it should be noted that although Gerngross discloses a genetically engineered microorganism expressing an exogenous α-1,2-mannosidase having an optimal pH between 5,1 and 8.0, however this is only a preferred process or a preferred embodiment of the teachings of Gerngross, particularly if one desires to have an "efficient" production of Man₅GlcNac₂ in vivo (see paragraphs 42, 68-72 and Table 6). The Gerngross reference does not teach explicitly that *T. reesei* α-1,2-mannosidase should not be used even though the reference acknowledges that this mannosidase as well as the *Aspergillus saitoi* α-1,2-mannosidase having a pH optimum around pH 5.0 are capable of producing Man₅GlcNac₂ in vivo even though they are not efficient (see at least paragraph 71). Moreover, Gemgross also teaches the use of exogenous enzymes selected either to have optimal activity under the conditions present in the fungi where

activity is desired or which are targeted to an organelle where optimal activity is achieved, such as a HDEL-T. reesei α -1,2-mannosidase fusion protein targeted to ER in a *Pichia pastoris* strain. This chimeric T. reesei α -1,2-mannosidase fusion protein would have intracellular enzymatic activity whereas the native T. reesei α -1,2-mannosidase does not have any intracellular enzymatic activity in a *Pichia pastoris* strain as evidenced at least by the teachings of Martinet et al. (Biotechnology Letters 20:1171-1177, 1998; IDS) discussed below.

Secondly, the rejected claims do not require the genetically engineered *Pichia* yeast strain to produce Man₅GicNAc₂ to any degree of efficiency or any particular fraction of produced glycoproteins having the structure Man₅GicNAc₂. Please also note that the term "reduced glycosylation" as defined by the instant specification to mean a reduced size of the carbohydrate moiety on the glycoprotein, particularly with fewer mannose residues, when the glycoproteinis expressed in a methylotrophic yeast strain which has been modified in accordance with the present invention, as compared to a wild type, unmodified strain of the methylotrophic yeast (page 28, lines 1-5).

2. With respect to the teaching of Gerngross relating to Och1, Gerngross merely discloses reducing endogenous mannosyltransferase activity and based on the disclosure of Gerngross it is unclear whether the disruption of OCH1 alone in a *Pichia* strain would be sufficient or whether multiple enzymes need to be disrupted in order to obtain Man₈ and ultimately Man₅ N-glycans. Although Gerngross refers to JP 8-336387 for an Och1 mutant strain of *P. pastoris*, this Japanese application does not appear to

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provide any showing that the predominant N-glycan form in the mutant is a Man₈ glycan. Applicants further argue that a recent report by Choi et al (Exhibit 1) showed that Man₉ is still a predominant N-glycan form in the Och1 mutant (Figure 3B).. Accordingly, Gerngross merely provides numerous potential options for those skilled in the art to experiment, and does not provide clear teaching that anticipates the presently claimed invention.

Firstly, with respect to Applicants' argument on whether multiple enzymes need to be disrupted in order to obtain Man₈ and ultimately Man₈ N-glycans, it is irrelevant because the claims do not require the genetically engineered yeast *Pichia* strain only having the disruption of OCH1 alone, and due to the open language of the term "comprising" in method claims.

Secondly, Figure 3 B in the post-filing art of Choi et al (Exhibit 1) only showed the released N-linked glycans analysis from K3 produced in och1 deleted *P.pastoris* strain BK64-1, not from och1 deleted *P. pastoris* strain expressing an exogenous α-1,2-mannosidase. Furthermore, Choi et al stated "Fungal α-1,2-mannosidase with acidic pH optima (e.g., *P citrinium* and *A. nidulans*), when expressed as fusions with the leader library, generally resulted in low (Man)₆(GlcNAc)₂ yields (data not shown) consistent with previous findings (11, 21)" (page 5026, col. 1, bottom of second full paragraph).

Accordingly, new claims 90-96 and 104-113 are still rejected under 35 U.S.C. 102(e) as being anticipated by Gerngross (US 2002/0137134; IDS) as evidenced by JP 8-336387 (IDS) for the reasons set forth above.

Claim Rejections - 35 USC § 103

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

This application currently names joint inventors. In considering patentability of the claims under 35 U.S.C. 103(a), the examiner presumes that the subject matter of the various claims was commonly owned at the time any inventions covered therein were made absent any evidence to the contrary. Applicant is advised of the obligation under 37 CFR 1.56 to point out the inventor and invention dates of each claim that was not commonly owned at the time a later invention was made in order for the examiner to consider the applicability of 35 U.S.C. 103(c) and potential 35 U.S.C. 102(e), (f) or (g) prior art under 35 U.S.C. 103(a).

New claims 90-91, 93-96, 104-108, 110 and 112-113 are rejected under 35 U.S.C. 103(a) as being unpatentable over Martinet et al. (Biotechnology Letters 20:1171-1177, 1998; IDS) in view of JP 8336387 (12/24/96; IDS) as evidenced by Choi et al. (PNAS 100:5022-5027, 2003; Cited by Applicants). This is a modified rejection necessitated by Applicant's amendment.

Martinet et al. teaches the preparation of plasmids for expression of *T. reesei* α-1.2-mannosidase or a chimeric *S. cerevisiae/ T. reesei* α-1.2-mannosidase (a fusion of

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the catalytic domain of *T. reesei* α-1,2-mannosidase to the ER retention signal of *S.* cerevisiae MNS1) in Pichia pastoris strains GSIV-HAs and GSIVNAf1s derived from the parental strain GS115 (see Materials and Methods, particularly sections "Strains and culture conditions" and "Construction of plasmids for expression of T. reesei α-1,2mannosidase in P. patoris"). Martinet et al. further teaches that in all expression plasmids are derived from the pPICZB vector, and the T. reesei α -1.2-mannosidase gene was under transcriptional control of the AOX1 promoter (page 1172, col. 2, first full paragraph). Martinet et al. also discloses that co-expression of heterologous T. reesei α-1,2-mannosidase in GSIVNAf1s resulting in partial trimming of the large influenza neuramidase (NA) N-glycans (>Man14GlcNac2) (see section "In vivo trimming of Nglycans by heterologous T. reesei α-1,2-mannosidase", and Figures 2A, 3). The coexpression of the chimeric MNS1/T. reesei α-1,2-mannosidase in GSIV-HAs resulted in the formation of both trimmed and hyperglycosylation glycan products of hemagglutinin (HA) (see page 1175, col. 2 and Fig. 4). Additionally, Martinet et al. notes that hyperglycosylation can be prevented by expression the protein of interest in the mutant veast strains mnn9, och1 or in the temperature-sensitive strain nad-29, where Nglycosylation is confined to the core oligosaccharide residues (page 1176, col. 1); and the results from the co-expression of the chimeric MNS1/T. reesei α-1,2-mannosidase in GSIV-Has suggest that removal of mannose residues creates more ideal substrates for P. pastoris mannosyltransferrases, leading to elongation of truncated glycosyl chains, and not to complete α -1,2-mannosidase digestion (page 1175, bottom of col. 2).

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Martinet et al. does not teach to further transform the *Pichia pastoris* strain with a vector comprising a portion of the Och1 gene and a selectable marker gene to effect the disruption of the genomic Och1 gene in the *Pichia pastoris* strain to reduce the glycosylation of a heterologous glycoprotein or producing a glycoprotein with reduced glycosylation; and a kit comprising the vectors and/or these further genetically modified *Pichia pastoris* yeasts.

However, at the effective filing date of the present application, JP 8-336387 already taught the preparation of a vector construct comprising a portion of *Pichia* OCH1 gene and a selectable marker gene for disruption of the genomic OCH1 in a *Pichia* yeast strain, including the <u>GTS 115 (NRRL Y-15851) strain</u> for inhibiting the elongation of sugar chains on glycoproteins for production of a glycoprotein having a sugar chain identical or similar to that of a medically useful biologically active protein (see at least the abstract in English, Fig. 9 on page 635 as well as col. 15, paragraph 33).

Accordingly, it would have been obvious and within the scope of skill for an ordinary artisan to modify the method and compositions taught by Martinet et al. by at least further transforming *Pichia pastoris* strains GSIV-HAs and GSIVNAf1s expressing heterologous T. reesei α -1,2-mannosidase using a vector construct comprising a portion of the *Pichia* OCH1 gene and a selectable marker gene for disruption of the genomic OCH1 taught by JP 8336387.

An ordinary skilled artisan would have been motivated to carry out the above modification because the elimination of endogenous OCH1 in a *Pichia* yeast strain

inhibits the addition of α-1,6-polymannose outer chain formation on the Asn-linked inner core oligosaccharide Man8GlcNAc2, and results in smaller and homogenous oligosaccharides in heterologous glycoproteins or at least production of a glycoprotein having a sugar chain identical or similar to that of a medically useful biologically active protein as taught by JP 8-336387. Moreover, Martinet et al. already noted at least that hyperglycosylation can be prevented by expression the protein of interest in the mutant yeast strains mnn9, och1 or in the temperature-sensitive strain ngd-29, where Nglycosylation is confined to the core oligosaccharide residues (page 1176, col. 1). The kits comprising the vector components for carrying out the modified methods and genetically modified Pichia pastoris yeast strains discussed above would also have been obvious. Furthermore, the genetically modified Pichia pastoris yeast strains resulting from the combined teachings of Martinet et al. and JP 8-336387 are capable of producing Man₅GlcNAc₂ as evidenced at least by the teachings of Choi et al which disclose fungal α-1,2-mannosidase with acidic pH optima (e.g., P citrinium and A. nidulans), when expressed as fusions with the leader library, in a P. pastoris och1 mutant strain generally resulted in low (Man)₅(GlcNAc)₂ yields consistent with previous findings (page 5026, col. 1, second full paragraph).

An ordinary skilled artisan would have a reasonable expectation of success to carry out the above modification in light of the teachings of Martinet et al., and JP 8-336387, coupled with a high level of skills of an ordinary skilled artisan in the relevant art.

Therefore, the claimed invention as a whole was prima facie obvious in the absence of evidence to the contrary.

Response to Arguments

Applicants' arguments related to the above rejection in the Amendment filed on 5/3/07 (pages 17-18) have been fully considered but they are respectfully not found persuasive for the reasons discussed below.

Applicants argue basically that Martinet et al reference refers to the S. cerevisiae mutant strains for preventing hyperglycosylation as evidenced by Exhibit 2, and that the reference also notes that the glycosylation pathways of S. cerevisiae and P. pastoris are significantly different. Applicants further argue that Man₉ is still a predominant N-glycan form in the Och1 mutant of Pichia pastoris as taught by Choi et al (Exhibit 1); and a skilled artisan would have expected to produce the Mans glycan structure from a Mans glycan, not from Man₉. Therefore, it would not have been obvious for an ordinary skilled artisan based on the disclosure of Martinet et al and JP 8336387 to obtain a Pichia pastoris strain as presently claimed and that is capable of producing the desired Mans glycan structure.

Firstly, it appears that Applicants considered each of the cited references in total isolation one from the other. JP 8-336387 already taught the preparation of a vector construct comprising a portion of Pichia OCH1 gene and a selectable marker gene for disruption of the genomic OCH1 in a Pichia yeast strain, including the GTS 115 (NRRL Y-15851) strain for inhibiting the elongation of sugar chains on glycoproteins for

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production of a glycoprotein having a sugar chain identical or similar to that of a medically useful biologically active protein. Particularly, the primary Martinet et al reference already notes that hyperglycosylation can be prevented by expression the protein of interest in the mutant yeast strains mnn9, och1 or in the temperature-sensitive strain ngd-29, where N-glycosylation is confined to the core oligosaccharide residues (page 1176, col. 1); and the results from the co-expression of the chimeric MNS1/T. reesei α -1,2-mannosidase in GSIV-Has suggest that removal of mannose residues creates more ideal substrates for P. pastoris mannosyltransferrases, leading to elongation of truncated glycosyl chains, and not to complete α -1,2-mannosidase digestion (page 1175, bottom of col. 2). Although the glycosylation pathways of S. cerevisiae and P. pastoris are significantly different, inactivated mutations including deletion of endogenous Och1 in either S. cerevisiae or P. pastoris strain would prevent or reduce hyperglycosylation.

Secondly, Figure 3 B in the post-filing art of Choi et al (Exhibit 1) only showed the released N-linked glycans analysis from K3 produced in och1 deleted P.pastoris strain BK64-1, not from och1 deleted P.pastoris strain expressing an exogenous α -1,2-mannosidase. Furthermore, Choi et al stated "Fungal α -1,2-mannosidase with acidic pH optima (e.g., P citrinium and A. nidulans), when expressed as fusions with the leader library, generally resulted in low (Man)₅(GlcNAc)₂ yields (data not shown) consistent with previous findings (11, 21)" (page 5026, col. 1, bottom of second full paragraph).

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Thirdly, the rejected claims do not require the genetically engineered *Pichia* yeast strain to produce Man₈GlcNAc₂ to any degree of efficiency or any particular fraction of produced glycoproteins having the structure Man₈GlcNAc₂. Please also note that the term "reduced glycosylation" as defined by the instant specification to mean a reduced size of the carbohydrate moiety on the glycoprotein, particularly with fewer mannose residues, when the glycoproteinis expressed in a methylotrophic yeast strain which has been modified in accordance with the present invention, as compared to a wild type, unmodified strain of the methylotrophic yeast (page 28, lines 1-5).

Accordingly, new claims 90-91, 93-96, 104-108 and110-113 are rejected under 35 U.S.C. 103(a) as being unpatentable over Martinet et al. (Biotechnology Letters 20:1171-1177, 1998; IDS) in view of JP 8336387 (12/24/96; IDS) as evidenced by Choi et al. (PNAS 100:5022-5027, 2003; Cited by Applicants) for the reasons set forth above.

In light of the Supreme Court Decision in KSR International Co. v. Teleflex Inc., 550 U.S.—82 USPQ2d 1385 (2007), the following rejection is applied.

New claims 97-106 are rejected under 35 U.S.C. 103(a) as being unpatentable over Martinet et al. (Biotechnology Letters 20:1171-1177, 1998; IDS) in view of JP 8336387 (12/24/96; IDS) as evidenced by Choi et al. (PNAS 100:5022-5027, 2003; Cited by Applicants) as applied to claims 90-91, 93-96, 104-108, 110 and 112-113 above, and further in view of Trombetta et al. (J. Biol. Chem. 271:27509-27516, 1996;

IDS) and Chiba et al. (J. Biol. Chem. 273:26295-26304, 1998; IDS). This is a new ground of rejection necessitated by Applicant's amendment.

The combined teachings of Martinet et al and JP 8336387 were already discussed above. However, none of the cited references teaches specifically that the genetically modified yeast *Pichia* further transformed with a vector comprising a nucleotide sequence coding for a glucosidase II or a functional part thereof or the use of a GAP promoter for expressing either *T. reesei* α-1.2-mannosidase and/or glucosidase II.

However, at the effective filing date of the present application Trombetta et al. already disclosed <u>cDNA</u> sequences encoding endoplasmic reticulum glucosidase II derived from various sources; including a glucosidase II gene from <u>S. cerevisiae</u> (see at least the abstract and sections titled "Primary sequence of α Subunit" and "Identification of the <u>S. cerevisiae</u> functional homologue of mammalian glucosidase II catalytic subunit (α) " on pages 27511-27513). Trombetta et al further demonstrated that <u>S. cerevisiae</u> functional homologue of mammalian glucosidase II catalytic subunit α removes two α -1,3-linked Glc units after removal of the terminal α -1,2-linked Glc residue in the core oligosaccharide Glc₃Man₆GlcNAc2 (page 17514, col. 1, second paragraph).

Additionally, Chiba et al already taught the preparation of an expression vector encoding HDEL-tagged Aspergillus α-1,2-mannosidase for expression in various Saccharomyces cerevisiae strains, named pGAMH1 plasmid containing the GAP promoter (see abstract and the section "DNA constructs. Chiba et al. further taught that carboxypeptidase Y produced in the YS132-8B yeasts having disrupted OCH1, MNN1

and MNN4 genes and harboring pGAMH1 plasmid has trimmed sugar chains up to Man₅GlcNAc₂, instead of carboxypeptidase Y containing high mannose type sugar chains in wild type Saccharomyces cerevisiae (see Fig. 1, and page 26302, col. 1, first full paragraph).

Accordingly, it would have been obvious and within the scope of skill for an ordinary artisan to further modify the combined teachings of Marinet et al. and JP 8336387 by further transforming a modified och-1 mutant *Pichia pastoris* strain expressing an exogenous *T. reesei* α -1,2-mannosidase tagged with an ER-retention signal with a recombinant vector expressing an exogenous ER glucosidase II, as well as the use of the GAP promoter for the expression of either the exogenous glucosidase II or *T. reesei* α -1,2-mannosidase in light of the teachings of Trombetta et al and Chiba et al as discussed above.

An ordinary skilled artisan would have been motivated to carry out the above modifications because expressing an exogenous glucosidase II in the modified och-1 mutant *Pichia pastoris* strain expressing heterologous *T. reesei* α-1,2-mannosidase would enhance the removal of any unprocessing glucose residues, and thus enhancing the enzymatic activity of the heterologous *T. reesei* I-1,2-mannosidase to result in a more complete trimming of sugar chains in heterologous glycoproteins expressing in these further modified *Pichia* yeast strains. Particularly, Martinet et al. already suggested that unprocessed glucose residues or capping glucose residues may block α-1,2-mannosidase treatment, and that these may be responsible for incomplete trimming of NA oligosaccharides to Man5GlcNAc2 (page 1175, col. 2,

first paragraph; page 1176, col.1, bottom of the second paragraph). Additionally, GAP promoter has been successfully used for expressing HDEL-tagged Aspergillus α -1,2-mannosidase in yeasts as already demonstrated by Chiba et al for generating heterologous carboxypeptidase Y having trimmed sugar chains up to Man₅GlcNAc₂.

An ordinary skilled artisan would have a reasonable expectation of success to carry out the above modifications in light of the teachings of Martinet et al., JP 8-336387, Trombetta et al. and Chiba et al., coupled with a high level of skills of an ordinary skilled artisan in the relevant art.

Therefore, the claimed invention as a whole was *prima facie* obvious in the absence of evidence to the contrary.

Double Patenting

The nonstatutory double patenting rejection is based on a judicially created doctrine grounded in public policy (a policy reflected in the statute) so as to prevent the unjustified or improper timewise extension of the "right to exclude" granted by a patent and to prevent possible harassment by multiple assignees. See *In re Goodman*, 11 F.3d 1046, 29 USPQ2d 2010 (Fed. Cir. 1993); *In re Longi*, 759 F.2d 887, 225 USPQ 645 (Fed. Cir. 1985); *In re Van Ornum*, 686 F.2d 937, 214 USPQ 761 (CCPA 1982); *In re Vogel*, 422 F.2d 438, 164 USPQ 619 (CCPA 1970);and, *In re Thorington*, 418 F.2d 528, 163 USPQ 644 (CCPA 1969).

A timely filed terminal disclaimer in compliance with 37 CFR 1.321(c) may be used to overcome an actual or provisional rejection based on a nonstatutory double patenting ground provided the conflicting application or patent is shown to be commonly owned with this application. See 37 CFR 1.130(b).

Effective January 1, 1994, a registered attorney or agent of record may sign a teminal disclaimer. A terminal disclaimer signed by the assignee must fully comply with 37 CFR 3,73(b).

New claims 90-96 and 105 are rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claim 9 of U.S. Patent No. 7,252,933. *This is a new ground of rejection necessitated by Applicant's amendment.*

Although the conflicting claims are not identical, they are not patentably distinct from each other because a methylotrophic yeast strain of any one of claims 1, 2, 3, 4-5 or 6-7, wherein the expression of said *T.reseei* α -1,2-mannosidase in said strain is directed by a promoter, wherein said promoter is the promoter of a gene selected from the group consisting of AOXI, an AOXII, GAP, YPT1 and FLD in the issued US Patent 7,252,933 anticipates the claimed genus of a genetically engineered strain of *Pichia* and a kit comprising the same in the application being examined and, therefore, a patent to the genus would, necessarily, extend the rights of the species or sub-should the genus issue as a patent after the species of sub-genus.

New claims 90-96 and 105 are provisionally rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims 39, 41-59 of copending Application No. 10/713,970. *This is a new ground of rejection necessitated by Applicant's amendment.*

Although the conflicting claims are not identical, they are not patentably distinct from each other because a genetically engineered *Pichia* strain, wherein said strain is engineered to express (1) a *Trischoderma reesei* α -1,2-mannosidase or a functional part thereof. (2) an N-acetylolucosaminyltransferase I (GnTI) or a functional part thereof.

and (3) a beta-1,4-galactosyltransferase (GaIT) or a functional part thereof, and the genomic OCH1 gene of said strain is disrupted in the copending Application No. 10/713,970 anticipates the claimed genus of a genetically engineered strain of *Pichia* and a kit comprising the same in the application being examined and, therefore, a patent to the genus would, necessarily, extend the rights of the species or sub-should the genus issue as a patent after the species of sub-genus.

This is a <u>provisional</u> obviousness-type double patenting rejection because the conflicting claims have not in fact been patented.

New claims 90-113 are rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims 5-8, 13-14 and 17-28 of U.S. Patent No. 6,803,225. *This is a new ground of rejection necessitated by Applicant's amendment.*

The claims of the present application differ from the claim of the issued US Patent 6.803,225 in reciting specifically a genetically engineered Pichia yeast strain expressing *T. reesei* α-1,2-mannosidase, and a recited Markush group of specific promoters used to express *T. reesei* α-1,2-mannosidase and/or glucosidase II.

The claims of the present application can not be considered to be patentably distinct over claims 5-8, 13-14 and 17-28 of U.S. Patent No. 6,803,225 when there is a specific disclosed embodiment of the issued US patent that teaches the use of vectors coding for T. reesei α -1,2-mannosidase and its expression under the control of at least

AOX1 promoter (see all the examples). Accordingly, the claims of the issued US patent fall within the scope of claims 90-113 of the present application.

This is because it would have been obvious to an ordinary skilled artisan to modify the claims of the issued US patent by also using vectors coding for T. reesei α -1,2-mannosidase and its expression under the control of at least AOX1 promoter for making and using the genetically engineered *Pichia* yeast strain, that support the instant claims. An ordinary skilled artisan would have been motivated to do this because this embodiment is explicitly disclosed or taught in the issued US patent as a preferred embodiment.

Conclusion

No claim is allowed.

Applicant's amendment necessitated the new ground(s) of rejection presented in this Office action. Accordingly, THIS ACTION IS MADE FINAL. See MPEP § 706.07(a). Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of

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the advisory action. In no event, however, will the statutory period for reply expire later

than SIX MONTHS from the date of this final action.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Quang Nguyen, Ph.D., whose telephone number is (571) 272-0776.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's SPE, Joseph T. Woitach, Ph.D., may be reached at (571) 272-0739.

To aid in correlating any papers for this application, all further correspondence regarding this application should be directed to Group Art Unit 1633; Central Fax No. (571) 273-8300.

Any inquiry of a general nature or relating to the status of this application or proceeding should be directed to (571) 272-0547.

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/QUANG NGUYEN, Ph.D./ Primary Examiner, Art Unit 1633